Claims

- 1. A recombinant RNA molecule which can be translated at least partly in a target cell, comprising a noninfectious virus genome of Coxsackie virus group B, preferably serotype B3, and at least one foreign gene which causes a desired function in the target cell, for example within the framework of a gene therapy.
- 2. The RNA molecule of claim 1, characterized in that it is replication-competent in the target cell.
- 3. The RNA molecule of claim 1 or 2, characterized in that in the virus genome parts of its coding sequence have been replaced by the at least one foreign gene.
- 4. The RNA molecule of claim 3, characterized in that in the virus genome the sequences of its capsid proteins VP1-VP4 have been replaced.
- 5. The RNA molecule of claim 3 or 4, characterized in that in the virus genome the sequences of its protease 2A and/or 3C have been replaced or modified such that there is no cytotoxicity for the target cell.
- 6. The RNA molecule of any of claims 3 to 5, characterized in that in the virus genome the sequences of its helicase 2C have been replaced.

- 7. The RNA molecule of any of claims 3 to 6, characterized in that in the virus genome the sequences of its protein 2B have been replaced.
- 8. The use of the RNA molecule of any of claims 1 to 7 for generating a vector for gene therapy.
- 9. A recombinant, infectious virion which is derived from Coxsackie Virus group B, preferably serotype B3, and whose genome is the RNA molecule of any of claims 1 to 7.
- 10. The virion of claim 9, characterized in that it corresponds in its structural proteins to a Coxsackie virus group B, preferably serotype B3.
- 11. A method for transducing a foreign gene into a target cell, comprising the steps
 - providing an RNA molecule of any of claims 1 to 7 or a virion of claim 9 or 10, and
 - infecting the target cell with the virion or transferring the RNA molecule by transfection.
- 12. A vector plasmid having at least one DNA sequence which codes for the RNA molecule of any of claims 1 to 7 and having a promoter located in front of the DNA sequence.
- 13. A helper construct for complementing the coding sequences replaced in the RNA molecule of any of claims 1 to 7.

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- 14. The helper construct of claim 13, characterized in that it is a helper plasmid which codes for at least one of the replaced sequences in a translatable manner.
- 15. The helper construct of claim 13, characterized in that it is a viral vector which codes for at least one of the replaced sequences in a translatable manner.
- 16. The helper construct of claim 13, characterized in that it is a helper cell which has been transfected stably with helper DNA coding for at least one of the replaced sequences.
- 17. A method for generating the virion of claim 9 or 10, comprising the steps:
 - transfecting of host cells with the vector plasmid of claim 12, and
 - complementing the replaced sequences in the host cell by the helper construct of any of claims 13 to 15.
- 18. The method of claim 17, characterized in that the host cell is the helper cell of claim 16.
- 19. A method for generating the vector plasmid of claim 12, comprising the steps

- a) providing a cDNA coding for infectious Coxsackie Viruses subgroup B, preferably subgroup B3,
- b) cloning the cDNA into a plasmid in a transcribable manner,
- c) amplifying sequence sections of the plasmid with the aid of primers leading to an amplificate which codes for the noninfectious virus genome, and
- d) ligating the amplificate to a DNA sequence for the foreign gene.
- 20. A method of generating the helper construct of any of claims 13 to 16, comprising the steps:
 - ruses subgroup B, preferably B3,
 - b) cloning the cDNA into a plasmid in a transcribable manner, and
 - c) amplifying sequence sections of the plasmid with the aid of primers leading to an amplificate which codes for the replaced coding sequences.
- 21. A kit, with the vector plasmid of claim 13 and a helper construct of any of claims 13 to 16.

- 22. A DNA molecule having at least one sequence section coding for the RNA molecule of any of claims 1 to 7.
- 23. A kit with a DNA molecule of claim 22.

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- 24. A kit for carrying out the method of claim 19 or 20, with
 - a plasmid containing cloned cDNA for infectious Coxsackie Viruses subgroup B, preferably subgroup B3,
 and
 - the primers required for amplification.
- 25. A therapeutic composition with the RNA molecule of any of claims 1 to 7.
- 26. A therapeutic composition with the vector plasmid of claim 12.
- 27. A therapeutic composition with virions of claim 9 or claim 10.
- 28. A DNA construct which codes for an RNA molecule of any of claims 1 to 7 and which persists and transcribes in a target cell but preferably does not replicate in the latter.
- 29. A recombinant virus, preferably adeno- or retrovirus, which codes for a recombinant RNA molecule of any of claims 1 to 7 and, after infection, expresses it in a

target cell, leading to a cytoplasmic replicon which is produced continuously.

- 30. A therapeutic composition with a virus of claim 29.
- 31. The use of the RNA molecule of any of claims 1 to 7 or of the virion of claim 9 or 10 for generating recombinant viruses or virions, preferably having a DNA genome, the foreign gene coding for gene functions lacking in the DNA genome.
- 32. A method for generating recombinant DNA viruses or DNA virions whose DNA genome lacks particular gene functions, in which method the missing gene functions are provided via a recombinant vector system with RNA genome.